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(54) Title: LARGE AREA SUBMUCOSAL GRAFT CONSTRUCTS AND METHOD FOR MAKING THE SAME

(57) Abstract

Multi-laminar tissue graft constructs, pseudoisotropic graft constructs and a method of preparing such constructs are described. Multi-laminar tissue graft constructs are prepared by fusing partially overlapped strips of submucosa tissue into a unitary graft construct by compressing the tissue under dehydrating conditions. Pseudoisotropic tissue graft constructs are prepared by overlapping a first strip of submucosa tissue with at least two strips of submucosa tissue orientated so that the longitudinal axis of each strip forms an angle of about 180°/N with the longitudinal axes of at least two other strips of submucosal tissue.

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LARGE AREA SUBMUCOSAL GRAFT CONSTRUCTS
AND METHOD FOR MAKING THE SAME

5 Field of the Invention

This invention relates to tissue graft constructs useful in promoting regrowth and healing of damaged or diseased tissue structures. More particularly this invention is directed to submucosal tissue graft constructs formed from multiple strips of submucosal tissue of a warm-blooded vertebrate and a method for making said constructs.

10 Background and Summary of the Invention

It is known that compositions comprising the tunica submucosa delaminated from both the tunica muscularis and at least the luminal portion of the tunica mucosa of the intestine of warm-blooded vertebrates can be used as tissue graft materials. See, for example, U.S. Patent Nos. 4,902,508 and 5,281,422. The compositions described in those patents are characterized by excellent mechanical properties, including high compliance, a high burst pressure point, and an effective porosity index which allows such compositions to be used beneficially for vascular graft constructs and in connective tissue replacement applications. When used in such applications the submucosal graft constructs appear to serve as a matrix for the regrowth of the tissues replaced by the graft constructs. Significantly, too, in over 600 cross-species implants, submucosa-derived graft compositions have never been shown to elucidate a tissue graft rejection reaction.

30 Submucosa-derived matrices for use in accordance with the present invention are collagen based biodegradable matrices comprising highly conserved collagens, glycoproteins, proteoglycans, and glycosaminoglycans in their natural configuration and natural concentration. One extracellular collagenous matrix for use in this invention

is submucosal tissue of a warm-blooded vertebrate.

Submucosal tissue can be obtained from various sources, for example, intestinal tissue harvested from animals raised for meat production, including, pigs, cattle and sheep or other warm-blooded vertebrates. Vertebrate submucosal tissue is a plentiful by-product of commercial meat production operations and is thus a low cost tissue graft material.

One limitation of the submucosal graft constructs described in the above mentioned patents is that the size of the graft is restricted by the size of the source material from which the submucosal tissue is prepared. For example, the size of a submucosal tissue graft prepared from intestinal tissues is limited by the length and circumference of the source segments intestinal tissue. Yet several applications of submucosal tissue graft constructs, including hernia repair, skin graft, meningeal coverings, repair of gastroschisis (congenital stomach defects) and organ tissue replacement, often require larger sheets of graft material than can be prepared directly from natural sources.

Large sheets of submucosal tissue can be prepared from smaller segments of submucosal tissue through conventional techniques such as weaving, knitting or the use of adhesives. However, commercial implementation of such techniques are often impractical and expensive. Additionally the use of adhesives or chemical pretreatment to promote adhesion of the tissue strips can compromise the biotrophic properties of the submucosal grafts. Thus there is a need for an inexpensive, easily manufactured, large area submucosal tissue graft construct that retains its biotrophic properties.

In accordance with one embodiment of the present application large area submucosal tissue graft constructs are formed from multiple pieces of vertebrate submucosa-

derived matrices. Unitary sheets (i.e., single piece graft constructs) of submucosal tissue are prepared in accordance with the present invention by fusing multiple strips of submucosal tissue to each other to form a sheet of tissue having a surface area larger than any one of the component strips of submucosal tissue. The present process comprises the steps of overlapping at least a portion of one strip of submucosal tissue with at least a portion of another strip of submucosal tissue and applying pressure at least to said overlapped portions under conditions allowing dehydration of the submucosal tissue. Under these conditions the overlapped portions will become "fused" to form a unitary large sheet of tissue. The large area graft constructs formed in accordance with the present invention consist essentially of submucosal tissue, free of potentially compromising adhesives and chemical pretreatments, and have a greater surface area and greater mechanical strength than the individual strips used to form the graft construct.

Individual strips of submucosal tissue as prepared from the tissues of a warm-blooded vertebrate have mechanical properties that are directionally specific (i.e. physical properties vary along different axes of the tissue). These directional characteristics are governed primarily by collagen orientation within the tissue. The collagen fibers are the load bearing constituents within intestinal submucosal tissue and are predominantly orientated parallel to the axis of the intestine lumen. This longitudinal disbursement of collagen in intestinal submucosal tissue contributes to the directional variability in physical properties of the submucosal tissue constructs.

In accordance with one embodiment of the present invention a unitary pseudoisotropic multi-laminate graft construct is prepared from multiple strips of submucosal tissue. The term "pseudoisotropic" as used herein

describes a graft material having approximately similar physical properties along each axis of the graft material. The pseudoisotropic multi-laminate graft constructs of the present invention are prepared from individual strips of submucosal tissue or sheets of submucosal tissue comprising strips of submucosal tissue. The method of preparing the pseudoisotropic graft constructs comprises overlapping a portion of a first strip (or sheet) with a second strip (or sheet), wherein the second strip (or sheet) is orientated in a plane parallel to the first strip (or sheet) but rotated so that the longitudinal axis of the first strip (or sheet) forms an angle relative to longitudinal axis of the second strip (or sheet). Additional strips (or sheets) can be added in a similar manner to create a multi-laminate structure having a desired number of laminate layers. The individual submucosal strips (or sheets) are then fused to one another to form a unitary multi-laminate pseudoisotropic construct by applying pressure at least to the overlapped portions of submucosal tissue.

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Brief Description of the Drawings

Fig. 1 is a diagrammatic representation of a homolaminate graft construct formed from multiple strips of submucosal tissue;

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Fig. 2 is a diagrammatic representation of a pseudoisotropic heterolaminate graft construct formed from four strips of submucosal tissue;

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Fig. 3 is a diagrammatic representation of a pseudoisotropic heterolaminate graft construct formed from three sheets of submucosal tissue, wherein each sheet is formed from multiple strips of submucosal tissue; and

Fig. 4 is a diagrammatic representation of one device suitable for forming perforations in submucosal tissue grafts in accordance with the present invention.

Detailed Description of the Preferred Embodiments

There is provided in accordance with this invention a method for forming tissue graft constructs comprising large area sheets of submucosal tissue. Furthermore, the present invention provides a method for preparing multi-laminate pseudoisotropic sheets of submucosal tissue. The method comprises the steps of fusing multiple strips of submucosal tissue to form unitary multi-laminar sheets of submucosal tissue.

The submucosal tissue suitable for use in the formation of the present graft constructs comprises naturally associated extracellular matrix proteins, glycoproteins and other factors. One source of submucosal tissue is the intestinal tissue of a warm-blooded vertebrate. Small intestinal tissue is a preferred source of submucosal tissue for use in this invention.

Suitable intestinal submucosal tissue typically comprises the tunica submucosa delaminated from both the tunica muscularis and at least the luminal portion of the tunica mucosa. In one embodiment of the present invention the intestinal submucosal tissue comprises the tunica submucosa and basilar portions of the tunica mucosa including the lamina muscularis mucosa and the stratum compactum which layers are known to vary in thickness and in definition dependent on the source vertebrate species.

The preparation of submucosal tissue for use in accordance with this invention is described in U.S. Patent No. 4,902,508, the disclosure of which is expressly incorporated herein by reference. A segment of vertebrate intestine, preferably harvested from porcine, ovine or bovine species, but not excluding other species, is subjected to abrasion using a longitudinal wiping motion to remove the outer layers, comprising smooth muscle tissues, and the innermost layer, i.e., the luminal portion of the

tunica mucosa. The submucosal tissue is rinsed with saline and optionally sterilized.

The multi-laminate submucosal tissue graft constructs of the present invention can be sterilized using conventional sterilization techniques including glutaraldehyde tanning, formaldehyde tanning at acidic pH, propylene oxide or ethylene oxide treatment, gas plasma sterilization, gamma radiation, electron beam, peracetic acid sterilization. Sterilization techniques which do not adversely affect the mechanical strength, structure, and biotropic properties of the submucosal tissue is preferred. For instance, strong gamma radiation may cause loss of strength of the sheets of submucosal tissue. Preferred sterilization techniques include exposing the graft to peracetic acid, 1-4 Mrads gamma irradiation (more preferably 1-2.5 Mrads of gamma irradiation), ethylene oxide treatment or gas plasma sterilization; peracetic acid sterilization is the most preferred sterilization method. Typically, the submucosal tissue is subjected to two or more sterilization processes. After the submucosal tissue is sterilized, for example by chemical treatment, the tissue may be wrapped in a plastic or foil wrap and sterilized again using electron beam or gamma irradiation sterilization techniques.

Submucosal tissue can be stored in a hydrated or dehydrated state. Lyophilized or air dried submucosa tissue can be rehydrated and used in accordance with this invention without significant loss of its biotropic and mechanical properties.

In one embodiment of this invention, large area compliant sheets of submucosal tissue are formed from multiple strips of submucosal tissue. The dimensions of the individual strips of submucosal tissue used is not critical and the term "strip of submucosal tissue" is defined herein to include submucosal tissue from one or

more vertebrate sources or organs in a wide variety of sizes and shapes. In one embodiment the strips are formed from a delaminated segment of intestinal tissue that is optionally but preferably cut and flattened out to provide 5 an elongated strips of submucosal tissue having two generally parallel sides and opposite ends. The term "sheet of submucosal tissue" is defined herein to include tissue constructs comprising multiple strips of submucosal tissue, wherein the strips are overlapped to form a 10 construct having a greater surface area than any one of the individual sheets used to form said construct. The term "layers of submucosal tissue" refers to the individual laminae of a multi-laminate submucosal tissue construct.

Unitary large area sheets of submucosal tissue 15 are formed in accordance with the present invention by overlapping individual strips of submucosal tissue and applying pressure to the overlapped portions to fuse the tissues together. In one embodiment pressure is applied to the overlapped tissue under conditions allowing dehydration 20 of the submucosal tissue. The large area sheets of submucosal tissue can be formed as either a heterolaminar sheet or a homolaminar sheet. The term "heterolaminar" as used herein refers to a multi-laminate tissue having a variable number of laminae of submucosa superimposed at 25 (and fused) at different points on the unitary graft construct. The term "homolaminar" as used herein refers to a multi-laminate tissue graft construct having a uniform number of laminae of submucosa at all points on the unitary graft construct.

In one embodiment the method of forming large 30 sheets of submucosal tissue comprises the steps of overlapping at least a portion of one strip of submucosal tissue with at least a portion of a second strip of submucosal tissue, and applying pressure at least to said 35 overlapped portions under conditions allowing dehydration

of the submucosal tissue. The amount of tissue overlap between the adjacent strips of submucosal tissue can be varied based on the intended use and the desired properties of the large area graft construct, provided that at least a portion of each strip of submucosal tissue overlaps with a portion of another strip of submucosal tissue. The applied pressure fuses the strips of submucosal tissue to one another along the overlapped portions, producing a compliant unitary heterolaminar sheet of submucosal tissue.

In another embodiment, a unitary homolaminate sheet of submucosal tissue can be prepared from strips of submucosal tissue. The method for forming the homolaminar tissue graft construct comprises the steps of forming a first layer of submucosal tissue, wherein strips of submucosal tissue are located side-by-side on a first surface. The strips of submucosal tissue of the first layer are located adjacent to one another so that the edges of the individual strips are in contact with one another without substantial overlap between one another. The first layer of submucosal tissue is then overlaid with a second layer of submucosal tissue. The strips of submucosal tissue of the second layer are located adjacent to one another similar to the strips of submucosal tissue of the first layer (i.e., adjacent to one another so that the edges of the individual strips are in contact with one another without substantial overlap between one another). In one embodiment the strips of submucosal tissue of the second layer are orientated in the same direction as the strips of submucosal tissue of the first layer, but offset in relationship to the submucosal strips of the first layer, so that the contacting edges of the individual strips of submucosal tissue of the first layer are bridged by the strips of submucosal tissue of the second layer (See Fig. 1). The overlap portions of the strips of submucosal tissue are then compressed between two surfaces, at least

one of the two surfaces being water permeable, under conditions allowing at least partial dehydration of the compressed submucosal tissue.

Advantageously, both the heterolaminar and homolaminar large area sheets of submucosal tissue formed in accordance with the present invention, consist essentially of submucosal tissue, have enhanced mechanical strength and have a greater surface area than any one of the individual strips used to form the submucosal sheets.

Submucosal tissue typically has an abluminal and a luminal surface. The luminal surface is the submucosal surface facing the lumen of the organ source and typically adjacent to an inner mucosa layer *in vivo* whereas the abluminal surface is the submucosal surface facing away from the lumen of the organ source and typically in contact with smooth muscle tissue *in vivo*. The multiple strips of submucosal tissue can be overlapped with the abluminal surface contacting the luminal surface, the luminal surface contacting the luminal surface or with the abluminal surface contacting the abluminal surface of an adjacent strip of submucosal tissue. All of these combinations of overlapping strips of submucosal tissue from the same or different vertebrate or organ sources will produce a large area sheet of submucosal tissue in accordance with the present invention upon compression of at least the overlapped portions under conditions allowing dehydration of the tissue.

The strips of submucosal tissue of the present invention can be conditioned, as described in U.S. Patent No. 5,275,826 (the disclosure of which is expressly incorporated herein by reference) to alter the viscoelastic properties of the submucosal tissue. In accordance with one embodiment submucosa delaminated from the tunica muscularis and luminal portion of the tunica mucosa is conditioned to have a strain of no more than 20%. The

submucosal tissue is conditioned by stretching, chemically treating, enzymatically treating or exposing the tissue to other environmental factors. In one embodiment the strips of intestinal submucosa tissue are conditioned by stretching in a longitudinal or lateral direction so that the strips of intestinal submucosa tissue have a strain of no more than 20%. The conditioned submucosal strips can be used to form large area sheets or multi-laminate structures in accordance with the present invention. Alternatively, the submucosal material can be conditioned after the formation of the large area sheet or multi-laminate large area sheet constructs to produce submucosal tissue material having a strain of no more than 20%.

During formation of the large area sheets of submucosal tissue, pressure is applied to the overlapped portions by compressing the submucosal tissue between two surfaces. The two surfaces can be formed from a variety of materials and in any shape depending on the desired form and specification of the unitary graft construct.

Typically the two surfaces are formed as flat plates but they can also include other shapes such as screens, opposed cylinders or rollers and complementary nonplanar surfaces. Each of these surfaces can optionally be heated or perforated. In preferred embodiments at least one of the two surfaces is water permeable. The term water permeable surface as used herein includes surfaces that are water absorbent, microporous or macroporous. Macroporous materials include perforated plates or meshes made of plastic, metal, ceramics or wood.

The submucosal tissue is compressed in accordance with one embodiment by placing the overlapped portions of the strips of submucosal tissue on a first surface and placing a second surface on top of the exposed submucosal surface. A force is then applied to bias the two surfaces towards one another, compressing the submucosal tissue

between the two surfaces. The biasing force can be generated by any number of methods known to those skilled in the art including the passage of the apparatus through a pair of pinch rollers (the distance between the surface of the two rollers being less than the original distance between the two plates), the application of a weight on the top plate, and the use of a hydraulic press or the application of atmospheric pressure on the two surfaces.

In one preferred embodiment the strips of submucosal tissue are subjected to conditions allowing dehydrating of the submucosal tissue concurrent with the compression of the tissue. The term "conditions allowing dehydration of the submucosal tissue" is defined to include any mechanical or environmental condition which promotes or induces the removal of water from the submucosal tissue at least at the points of overlap. To promote dehydration of the compressed submucosal tissue, at least one of the two surfaces compressing the tissue may be water permeable. Dehydration of the tissue can optionally be further enhanced by applying blotting material, heating the tissue or blowing air across the exterior of the two compressing surfaces.

The multiple strips of submucosal tissue are typically compressed for 12-48 hours at room temperature, although heat may also be applied. For example a warming blanket can be applied to the exterior of the compressing surfaces to raise the temperature of the compressed tissue up to about 40°C to about 50°C. The overlapped portions are usually compressed for a length of time determined by the degree of dehydration of the tissue. The use of heat increases the rate of dehydration and thus decreases the amount of time the overlapped portions of tissue are required to be compressed. Typically the tissue is compressed for a sufficient time to produce a stiff but flexible material. Sufficient dehydration of the tissue

is also indicated by a increase in impedance of electrical current flowing through the tissue. When impedance has increased by 100-200 ohms, the tissue is sufficiently dehydrated and the pressure can be released.

5 The compressed submucosal tissue can be removed from the two surfaces as a unitary compliant large area tissue construct. The construct can be further manipulated (i.e., cut, folded, sutured, etc.) to suit various medical applications where the submucosal material of the present invention is required.

10 A vacuum can optionally be applied to submucosal tissue during the compression procedure. The applied vacuum enhances the dehydration of the tissue and may assist the compression of the tissue. Alternatively the application of a vacuum may provide the sole compressing force for compressing the overlapped portions of the multiple strips of submucosal tissue. For example the overlapped submucosal tissue is laid out between two surfaces, preferable one of which is water permeable. The apparatus is covered with blotting material, to soak up water, and a breather blanket to allow air flow. The apparatus is then placed in a vacuum chamber and a vacuum is applied, generally ranging from 14-70 inches of Hg (7-35 psi). Preferably a vacuum is applied at approximately 51 inches of Hg (25 psi). Optionally a heating blanket can be placed on top of the chamber to heat the submucosal tissue during the compression of the tissue. Chambers suitable for use in this embodiment are known to those skilled in the art and include any device that is equipped with a vacuum port. The resulting drop in atmospheric pressure coacts with the two surfaces to compress the submucosal tissue and simultaneously dehydrate the submucosal tissue.

15 20 25 30 35 Optionally, large area tissue grafts can be formed into various shapes for tissue graft applications. For example, in organ reconstruction applications the large

area sheets can be formed in the shape of a hollow sphere or pouch. Such a shaped construct would be advantageous in the replacement of large regions of the urinary bladder or stomach. These shaped submucosal tissue constructs can be formed by conventional techniques such as cutting and suturing the tissue to form a desired shape.

Alternatively, strips of submucosal tissue can be formed into a large sheet of submucosal tissue having a nonplanar shape through a simple manufacturing procedure. The method comprises the steps of placing multiple strips of submucosal tissue between two complementary nonplanar shaped surfaces and compressing the overlapped strips of submucosal tissue between the two surfaces. The complementary shaped surfaces are formed such that the two surfaces can be pressed together such that the surfaces fit snug against one another without leaving any substantial pockets of air between the two surfaces. Preferably at least one of the two complementary surfaces is water permeable.

20 One method of forming a shaped submucosal construct comprises placing multiple strips of submucosal tissue on a nonplanar shaped porous surface such that the submucosal tissue conforms to the shape of the porous surface. Preferably the submucosal tissue is placed on the porous surface without stretching the material, however, the submucosal tissue can be stretched to facilitate covering the shaped porous surface. Each of the strips of submucosal tissue is positioned on the porous surface to overlap at least a portion of an adjacent strip of
25 submucosal tissue. The overlapped portions of the submucosal tissue are then covered with a second shaped surface that is complementary in shape with the first porous surface and pressure is applied to compress the submucosal tissue between the two surfaces under conditions
30 allowing dehydration of the submucosal tissue.

Alternatively the large area sheets of the present invention can be shaped into a nonplanar shape by stretching the large area sheet through the use of a die press procedure, wherein the submucosal tissue is pressed into a nonplanar shape by a porous die under dehydrating conditions such that the formed tissue graft holds its shape. Preferably a multi-laminate large area sheet is used in such a procedure.

Multi-laminar submucosal tissue constructs are formed in accordance with the present invention by overlapping a portion of one strip of submucosal tissue with a portion of another strip of submucosal tissue. In a similar fashion large area multi-laminar tissue graft constructs can be formed in accordance with the present invention by overlapping a sheet of submucosal tissue (formed as described above) with at least a portion of a second sheet of submucosal tissue. The size and physical properties of the multi-laminate submucosal tissue construct can be regulated by the number of overlapped strips of submucosal tissue and the percent of the overlapped portion of each strip.

The multi-laminar tissue graft constructs are formed in accordance with the present invention by overlapping a portion of one strip of submucosal tissue with a portion of another strip of submucosal tissue to form a first sheet. Additional strips of submucosal tissue are overlaid onto the overlapped portions of the first sheet to form a second sheet, wherein the edges of the strips of the second sheet are optionally at an acute angle to the edges of the strips in the first sheet, and wherein said formed second sheet is coplanar with the first sheet. The strips of submucosal tissue of the second sheet can be positioned so that at least a portion of one strip of submucosal tissue of the second sheet overlaps with at least a portion of another strip of submucosal tissue of

the second sheet. Additional strips of submucosal tissue can be overlaid on the overlapped portions of the first and second sheets to provide additional layers of submucosal tissue. The multiple layers of submucosal tissue are then compressed under dehydrating conditions to form a multi-ply heterolaminar submucosal tissue construct having a surface area greater than any one of the individual strips of submucosal tissue used to form the multilayered construct.

In one embodiment of the present invention submucosal tissue is cut into strips, each strip having generally parallel sides, and used to form the multilayered heterolaminar sheets of the present invention. In this embodiment the strips of submucosal tissue of the second layer are overlaid onto the overlapped portions of the first layer such that the edges of the first layer submucosal strips are at an angle relative to the edges of the second layer submucosal strips. The overlapped portions of submucosal tissue are compressed under dehydrating conditions to form the multilayered heterolaminar sheet. These sheets can be cut without unraveling and do not delaminate when soaked in water for a period of time (greater than one hour) that corresponds to the time required for implanting the sheet in a host. If the sheet is properly implanted such that it is sutured on all sides, delamination should not occur subsequent to implantation.

In accordance with the present invention a tissue graft construct having pseudoisotropic properties is formed from strips of submucosal tissue. The pseudoisotropic tissue graft constructs comprise multiple strips of submucosal tissue fused together, in the absence of adhesives or sutures, in a multi-laminate structure. The pseudoisotropic multi-laminate tissue graft is prepared in accordance with the present invention from at least three strips of intestinal submucosal tissue delaminated from

both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate. Each of the strips of intestinal submucosal tissue are characterized as having a longitudinal axis corresponding to the predominant orientation of the collagen fibers in the submucosal tissue strips. The method of forming the pseudoisotropic graft constructs comprises locating a first strip of submucosal tissue on a first surface, overlaying said first strip with at least two additional strips of submucosal tissue so that the longitudinal axes of each individual strip of submucosal tissue forms an angle of about $180^\circ/N$ with the longitudinal axis of at least two other strips of submucosal tissue forming the heterolaminate graft, wherein N = the total number of strips of submucosal tissue. (See Fig. 2). For example a pseudoisotropic graft construct formed from four (4) strips of submucosal tissue will have an angle of 45° ($180^\circ/4 = 45^\circ$) formed between the central longitudinal axes of each strip in reference to two of the other three strips forming the graft construct. (See Fig. 2). The submucosal tissue (at least the overlapped portions) is then compressed between the first surface and a second surface. In one embodiment the tissue is compressed under conditions allowing at least partial dehydration of the compressed submucosal tissue, and in a preferred embodiment at least one of said surfaces is water permeable.

Large area tissue graft constructs having pseudoisotropic properties can also be prepared from large area sheets of submucosal tissue. These pseudoisotropic tissue graft constructs comprise multiple layers of large area sheets of submucosal tissue wherein the sheets of submucosal tissue comprise overlapped strips of submucosal tissue. The sheets of submucosal tissue, comprising overlapped strips of submucosal tissue, can be formed as

homolaminar or heterolaminar tissue constructs as described previously in the present description (See Fig. 3).

One method of preparing a large area multi-laminate tissue graft construct having pseudoisotrophic properties comprises forming a first sheet of submucosal tissue from multiple strips of submucosal tissue and overlaying the first sheet with at least two additional sheets. The individual strips of submucosal tissue comprising each sheet have a longitudinal axis corresponding to the predominant orientation of the collagen fibers in the submucosal tissue strips. The first sheet is formed on a first surface by overlapping the individual strips of submucosal tissue so that each strip is aligned with the adjacent strips and the longitudinal axis of each strip of submucosal tissue are substantially parallel to one another. Thus the collagen fibers of the first sheet are aligned predominantly in a single orientation, such that the sheet can be characterized as having a longitudinal axis corresponding to the predominant orientation of the collagen fibers.

As described above, large area sheets of submucosal tissue can be formed from overlapped submucosal tissue to form either heterolaminar or homolaminar sheets of submucosal tissue. Both heterolaminar and homolaminar sheets are suitable for forming the large area pseudoisotrophic tissue graft constructs of the present invention.

After the first sheet of submucosal tissue is located on a first surface, additional layers of submucosal sheets are formed on top of the first sheet in the same manner that the first sheet was formed (i.e. each sheet of submucosal tissue of the multi-laminate comprises overlapped strips of submucosal tissue wherein the longitudinal axes of the strips of submucosal tissue comprising each sheet are substantially parallel to one

another). Each individual sheet is overlaid on another sheet so that the longitudinal axes of the strips of submucosal tissue of the overlaid sheet forms an angle of about $180^\circ/S$. (S = the total number of sheets of submucosal tissue) with the longitudinal axes of the strips of submucosal tissue of at least two of the other sheets forming the multi-laminate construct. Once the total number of sheets have been overlaid, the sheets of submucosal tissue are compressed between the first surface and a second surface under conditions allowing at least partial dehydration of the compressed submucosal tissue. In preferred embodiments at least one of said surfaces is water permeable.

In one embodiment, after multiple strips of submucosal tissue are overlapped with one another, the overlapped portions are manipulated to remove trapped air and bulk quantities of water before fusing the strips into a single sheet of submucosal tissue. In general the trapped air bubbles and bulk quantities of water are squeezed out through the use of a compressing force which is moved across the surface of the overlapped portions. The compressing force can take the form of a cylinder that is rolled across the surface of the overlapped portions, or alternatively the overlapped portions can be passed between two or more rollers wherein the distance between the surface of the opposing rollers is less than the thickness of the submucosal sheet. The overlapped portions can then be compressed if necessary for an additional length of time under dehydrating conditions to fuse the multiple strips into a single sheet of submucosal tissue in accordance with the present invention.

The excess portions of the pseudoisotropic multi-laminate grafts (i.e., those portions of the graft having a laminate number less than N or S) can be removed after formation of the multi-laminate. Furthermore, the

mechanical properties of multi-laminate submucosal material can be tailored to the medical application needs by adjusting the percentage of overlap between adjacent strips of submucosal tissue, altering the number of submucosal tissue layers, varying the angle of adjacent layers relative to one another, changing the water permeability of the compressing surfaces and/or the composition of the compressing surfaces, selecting the shape of the compressive surfaces, and varying the load applied to compress the overlapped submucosal tissue.

The pseudolaminate submucosa material of the present invention can be further modified to include a plurality of perforations. Recent experiments have demonstrated that the process of remodelling is slower with implanted multi-laminate submucosal tissue graft constructs than for single or two layered submucosal tissue grafts. In addition, multi-laminate submucosal tissue graft constructs tend to accumulate tissue fluid in cyst-like pockets between adjacent laminae during the first 14-28 days after implantation in soft tissue locations (such as the muscular body wall of rats). Fluid pockets are detrimental to wound healing because they retard connective tissue ingrowth, provide an environment conducive to bacterial growth, and prevent the apposition of natural (native) body tissues which promotes healing and tensile strength.

Perforation of multi-laminate graft constructs has been found to enhance the graft's *in vivo* remodelling properties and to enhance the adhesion of the tissue graft layers to one another. The perforations are believed to promote contact of the submucosal tissue with endogenous fluids and cells (by increasing the surface area of the implanted graft) and the perforations also serve as a conduit allowing extracellular fluid to pass through the graft.

In accordance with the present invention the term "perforate" designates a bore that extends through the entire graft construct. However, tissue graft constructs having "holes", defined herein as a cavity that penetrates into the tissue but does not extend through the entire graft construct are also within the scope of the present invention. The spacing and size of the perforations, as well as the depth to which the perforations penetrate into the tissue, will be varied according to the desired mechanical strength, porosity, size and thickness (number of layers) and other factors related to the medical application of the tissue graft. The size of the perforations range from 0.5 to 3 mm, more preferably from 0.6 to 2 mm. In one embodiment the perforations are even spaced from one another at a distance ranging from 2 to 20 mm, more preferably from 3 to 7 mm.

Typically the perforations are formed in the submucosal tissue while the tissue remains at least partially hydrated. In the pseudoisotropic multi-laminate submucosal tissue materials of the present invention, the perforations are preferably made after formation of the multi-laminate construct and when the tissue has been dried to a water content of approximately 10-20% by weight water (10-20% hydrated). Sufficient drying of the tissue can be determined by weighing the fresh tissue and drying the tissue to 10-20% of the fresh weight or the sufficient drying can be determined by impedance measurement as previously described. After perforation of the tissue the submucosal tissue is subjected to terminal sterilization and stored as described previously.

In one embodiment holes (extending only part way through the tissue) or perforations can be formed on both sides of the tissue graft. In addition the tissue can be modified to include perforations as well as holes that extend only part way through the tissue. Furthermore the

submucosal tissue can be modified to include a plurality of holes, wherein various subsets of holes extend to different depths into the tissue relative to the other formed holes. This can be accomplished, for example, by perforating the individual layers of submucosal tissue before overlapping the layers to form the multi-laminate construct. If some of the layers are not perforated or if the perforations of the individual layers are not aligned, the formed multi-laminate construct will have holes extending to different depths into the tissue. Preferably the tissue is perforated, in a uniform distribution over the surfaces of the tissue graft, thus forming a series of bores that allow fluid communication from a the first planar surface to a second opposite planar surface of the graft construct.

In one embodiment the perforations are formed perpendicular to the surface of the tissue graft construct, i.e., the longitudinal axis of the perforation/hole forms a 90° angle with the plane defining the surface of the graft. Alternatively the perforations can be formed so that the axis of perforation is not perpendicular to the surface of the graft (i.e. so that a longitudinal axis parallel to the wall defining the perforation/hole forms an angle other than 90° with the plane of the graft surface). In accordance with one embodiment the perforations are formed at an angle ranging from 45° to 90° in reference to the surface of the graft.

Multi-laminar tissue grafts can be cut without unraveling and do not delaminate when soaked in water for a period of time (greater than one hour) that corresponds to the time required for implanting the sheet in a host. However, multi-laminate tissue constructs tend to accumulate tissue fluid in cyst-like pockets between adjacent laminae during the first 14-28 days after implantation in soft tissue locations (such as the muscular body wall of rats). Perforations of the pseudoisotropic

multi-laminate graft constructs of the present invention will alleviate the accumulation of fluids between the layers of the multi-laminar construct by providing a conduit through which the fluid can flow out of the tissue.

5 In addition the perforations will have a "stapling" effect that will augment the adhesion of the laminae to each other.

10 Accordingly, the placement of full thickness or partial thickness holes in multi-laminate tissue grafts provide the following advantages over non-perforated multi-laminate sheets:

1. Increased passage of fluids (including tissue fluids) through the material; and
2. Increased adhesive force between adjacent layers.

15 The submucosal tissue can be perforated using a wide variety of devices known to those skilled in the art. The method utilized to perforate the submucosal tissue is not critical provided the aggregate structural integrity of the submucosal tissue is maintained.

20 In preferred embodiments the perforation of the submucosal tissue does not result in the removal of significant amounts of tissue. For example, the perforations are formed by pressing a pointed solid object through the tissue to press the tissue aside during the insertion of a solid object, as opposed to boring out the material. Other means for perforating the tissue include the use of ballistics, cutting instruments, laser beams or enzymatic or chemical treatments.

25 In one embodiment, the submucosal tissue is perforated by pressing a pin or solid needle, into/through the tissue. Typically a 20-23 gauge solid needle is used to form the perforations. In this manner, no significant amount tissue is removed during the process of forming the perforations, but rather a portion of each layer is torn and pushed into an adjacent layer to provide a stapling

effect. This "stapling" effect can be further enhanced by forming a portion of the perforations from one side of the graft and forming the remaining perforations from the opposite side of the graft.

Fig. 4 depicts one embodiment of a device for perforating the submucosal tissue graft constructs. The device comprises a base (1) and a plurality of stainless steel pins (2) embedded in base (1) and extending out through the surface of the base. The base comprises Epoxy (E) and Delrin (D) portions and has a length of 3.2 inches, a width of 1.85 inches and is 0.5 inches thick. The Epoxy and Delrin portions each have a length of 3.2 inches, a width of 1.85 inches and are 0.25 inches thick. In accordance with this embodiment pins (2) are substantially parallel to one another and form a 90° angle with the surface of the base. Pins (2) are 0.040 inches in diameter, are spaced 0.264 inches apart (center to center of adjacent pins) within a 0.4 border from the edge of the device and protrude 0.25 in. from the base. The device thus holds a total of fifty pins.

Example 1

Submucosal tissue was prepared from vertebrate intestinal tissue in accordance with the procedure described in U.S. Patent No. 4,902,508. Strips of submucosal tissue were formed from a segment of intestinal tissue of a warm-blooded vertebrate, said segment comprising the tunica submucosa delaminated from both the tunica muscularis and at least the luminal portion of the tunica mucosa of said segment of intestinal tissue. The segment of intestinal tissue was cut along the longitudinal axis of the segment and laid flat. The tissue was then further sliced into a series of strips each having generally parallel sides.

Multiple strips of submucosal tissue were organized on a 12 by 12 inch perforated stainless steel plate wherein a portion of one strip of submucosal tissue overlaps a portion of the adjacent strip of submucosal tissue. A second 12 by 12 inch perforated stainless steel plate was then placed on top of the submucosal tissue. The perforated stainless steel plates used in this embodiment has 0.045 inch perforations arranged straight center and located 0.066 inches apart. A 50-100 pound weight was placed on top of the second stainless steel plate and the tissue was compressed for 24 hours at room temperature.

Example 2

Strips of submucosal tissue were prepared as described in Example 1. Multiple strips of submucosal tissue were laid out between two perforated, stainless steel plates so that a portion of one strip of submucosal tissue overlapped a portion of the adjacent strip of submucosal tissue. The "plate-submucosa-plate" apparatus was placed on a flat surface and covered with blotting material, to soak up water, and a breather blanket to allow air flow. The apparatus was then sealed into a nylon bag that has a vacuum port. A vacuum was applied (greater than 28 inches of Hg) to pull air out of the vacuum bag and the resulting drop in atmospheric pressure simultaneously compressed and dehydrated the submucosal tissue. After 24 hours of applying a vacuum, the produced sheet was moist and very flexible. No seams from the layering of the submucosal tissue were visible and the strength of a prototype 8-thickness sheet as determined by ball burst test was 80 pounds.

Example 3

Strips of submucosal tissue were prepared as described in Example 1. The submucosal tissue strips were organized on a mesh so that a portion of one strip of submucosal tissue overlapped a portion of the adjacent strip of submucosal tissue. Once the mesh was covered with one layer of submucosal tissue a second layer of submucosal tissue was applied on top of the first layer so that the edges of the submucosal strips of the second layer were at an angle relative to edges of the submucosal strips of the first layer.

After all the strips of submucosal tissue were placed on the mesh, another mesh was placed on top of the submucosal tissue layers and the "mesh-submucosal tissue-mesh" sandwich was compressed with a load and dried. This process produced a dried large area submucosal sheet that was peeled off the mesh as a unitary graft construct.

Example 4Sterilization of Submucosal Tissue with Peracetic Acid

Submucosal tissue is soaked in a peracetic acid/ethanol solution for 2 hours at room temperature using a ratio of 20:1 (mls peracetic solution: grams submucosal tissue) or greater. The peracetic acid/ethanol solution comprises 4% ethanol, 0.1% (volume: volume) peracetic acid and the remainder water. The 0.1% peracetic acid component is a dilution of a 35% peracetic acid stock solution commercially available and defined as in table 1.

Preferably, the submucosal tissue is shaken on a rotator while soaking in the peracetic acid solution. After two hours, the peracetic acid solution is poured off and replaced with an equivalent amount of lactated Ringer's solution or phosphate buffered saline (PBS) and soaked

(with shaking) for 15 minutes. The submucosal tissue is subjected to four more cycles of washing with lactated Ringer's or PBS and then rinsed with sterile water for an additional 15 minutes.

5

Table 1: Chemical Composition of the 35% Peracetic Acid Solution

Composition, % by weight

10	Peracetic acid	35.5
	Hydrogen peroxide	6.8
	Acetic acid	39.3
	Sulfuric acid	1.0
	Water	17.4
15	Acetyl peroxide	0.0
	Stabilizer	500 PPM

Typical active oxygen analysis, % by weight

20	Active Oxygen as peracid	7.47
	Active Oxygen as H ₂ O ₂	2.40
	Total active oxygen	10.67

Sterilization of Submucosal Tissue with Ethylene Oxide

After preparation of the multi-laminate constructs using sterile conditions, the material is packaged and subjected to a second round of sterilization (terminal sterilization). The tissue can be packaged in plastic that is permeable to ethylene oxide and subjected to ethylene sterilization according to procedures known to those skilled in the art. Essentially the packaged material is exposed to ethylene oxide for four hours at 115° F. During the sterilization the tissue is also provided with 65% relative humidity for at least 75 minutes of the 4 hour treatment. The high humidity enhances uptake of the ethylene oxide by the tissue. After four hours the

30

35

ethylene oxide, ethylene chlorohydrone and ethylene glycol is flushed out with nitrogen and air.

Example 5

5

Ball Burst Strength Test by Means of a Compression Cage and a MTS Tensile Tester

The strength of multi-laminate submucosal tissue grafts is determined through the use of a material testing system (MTS) tensile tester. The multi-laminate tissue construct is secured within a four sided frame clamp (specimen clamp) to provide uniform distribution of the stress through out the tissue construct. The initial fixture level is set so that the top of the steel ball is located immediately under the plane the test specimen. The handle of the specimen clamp is lifted to its topmost position so that the jaws of the clamp are able to accept the test specimen. The submucosal tissue construct is cut to fit the specimen clamp, the aperture of the clamp having a diameter of one and three-quarter inches. A half-inch of excess material should be included around the perimeter of the test specimen to ensure sufficient clamping area. The submucosal tissue is placed in jaws of the clamp and secured, the clamp force being controlled by thumbwheel means located on the top clamp.

The clamped submucosal tissue is then pressed down over a metal ball at a controlled rate utilizing a tensile tester software interface to control and measure the force placed on the test specimen. The force is increased until failure of the specimen occurs. Failure is defined as the maximum load which corresponds to the first appearance of the ball through visible non-natural discontinuities in the plane of the specimen. In the case that the topmost position of the fixture is reached prior to failure, the software limits will engage and discontinue

30

35

the test. The peak load value displayed on the Microp profiler 458.01 is recorded and the specimen is removed.

5 Example 6

A multi-laminate tissue graft construct was prepared as follows:

An ample amount of submucosal tissue is prepared from vertebrate intestine, cut and flattened out and disinfected with peracetic acid as described in Example 4 (approximately 70 grams of submucosal tissue is required for a 10 cm x 15 cm device). Surgical gloves, face mask, and cap should be worn after sterilization of the tissue with peracetic acid to minimize the contamination from organic matter and airborne particulate.

Strips of submucosal tissue are placed on top of a first perforated stainless steel plate in the desired orientation. The stainless steel plates used are perforated stainless steel plates with 0.045 inch round perforations on straight centers and located 0.066 inches apart. After formation of a layer of submucosal tissue the submucosal tissue is smoothed out to remove air bubbles. Additional layers are overlaid until the device is complete. Excess material is removed from around the multi-laminate structure with a scissors. The weight of the submucosal multi-laminate is recorded. A second stainless steel plate (perforated with 0.045 inch round perforations on straight centers located 0.066 inches apart) is placed on top of the multi-laminate construct.

The multi-laminate construct can optionally be "pinch rolled" to remove trapped air and water. To pinch roll the material, the two perforated metal plates surrounding the submucosal tissue are placed in-between two polypropylene sheets (Kimberly Clark, class 100 "Crew

Wipe") and the entire apparatus is placed in-between two layers of nylon bagging film (Zip Vac, Auburn WA) that are larger than 1"x 1". A weighted cylinder is then rolled across the apparatus numerous times (at least three times).

5 To perforate the multi-laminate submucosal tissue graft construct the apparatus is partially disassembled to expose the top surface of the tissue graft and a piece of nylon bagging film is placed directly on the top most layer of submucosa tissue. The multi-laminate submucosal tissue graft construct is then inverted onto a Styrofoam work surface, and the first stainless steel plate is carefully removed. The exposed surface of submucosa is then covered with a piece of nylon bagging film. The tissue graft construct is then perforated, and then the top nylon
10 bagging film is removed. The multi-laminate submucosal tissue graft construct is then re-inverted and placed back on the perforated stainless steel plate. The nylon bagging film is removed from the submucosa top surface and a second perforated stainless steel plate is placed on top of the
15 multi-laminate submucosal tissue graft construct.
20

The multi-laminate submucosal tissue graft construct is then compressed under dehydrating conditions as follows:

A layer of blotting material (NuGauze) larger
25 than the size of the perforated plates is placed on a table top (or other smooth level surface). The stainless steel plates with the multi-laminate submucosal tissue graft construct between them is placed on top of the blotting material. Another layer of blotting material
30 (approximately the same size as the first sheet of blotting material) is placed on top of the stainless steel plates. A breather blanket (Zip Vac, Auburn, WA) is placed on top of the blotting material. Preferably the breather blanket is slightly larger than the objects it is covering.

Optionally electrodes can be placed in contact with the submucosal tissue to allow the measurement of impedance across the tissue. Typically the tissue is compressed for a sufficient time to produce a stiff but flexible material. Sufficient dehydration of the tissue is indicated by a increase in impedance of electrical current flowing through the tissue. When impedance has increased by 100-200 ohms, the tissue is sufficiently dehydrated and the pressure can be released.

A border of chromate tape is placed on the table top around the apparatus and the area to be vacuum pressed. The backing is removed from the tape and a piece of the nylon bagging film that already has the nozzle port attached to it is placed on top of the area enclosed by die chromate tape (see Figures 3a & 3b) and adhered to the tape. The heating blanket, if used is turned on, and vacuum pump is turned on. The bag should be checked for wrinkles (smooth them out if found) and for an inadequate seal between the chromate tape and the nylon bagging film (correct if found). A vacuum should be drawn to a level ranging from 25 to 30 psi_{vac}. After vacuuming to the desired hydration level (approximately 24 hours), the seal of the bag is broken at a taped region, the vacuum pump is turned off and the unitary, perforated multi-laminate submucosal tissue graft construct is removed. A pair of scissors can be used to cut off any portion of the tissue graft that did not receive the complete amount of overlap.

Example 7

The submucosal tissue graft construct can also be perforated after formation of the unitary multi-laminate construct as follows. The multi-laminate construct is formed in accordance with Example 3. The mesh/submucosal tissue sandwich was removed from the drying apparatus and

the tissue was perforated. The graft was perforated by inserting a nail between the mesh of the wire and pushing the nail through the tissue at multiple points on the graft surface. The perforated multi-laminate submucosal tissue was then cut square (4 1/2 x 4 1/2 in.) and marked for identification purposes.

5 **Example 8**

10 A perforated pseudoisolaminate construct was prepared as follows: Strips of submucosal tissue were arranged in 4 layers on a wire mesh. The first layer was laid directly on the mesh and the remaining three layers were overlaid on top of the first layer at an angles of 45°, 15 90° and 135° relative to the first layer, respectively (see Fig. 2). A second mesh was placed on top of the submucosal tissue, and the tissue was sandwiched between the mesh and c-clamped to the drying rack. A fan was placed in front of the rack and turned on. Holes were punched through the 20 tissue in a checkerboard pattern using the mesh as a guide. (i.e., every alternate space in the mesh was used to perforate the tissue. Accordingly the pattern appeared as follows:

25 X X X
 X X
 X X X
 X X

30 The perforation of the tissue was stopped before completion because the submucosal tissue was being disrupted. Therefore the submucosal tissue was dried for 25 min. with the fan on high. The remaining perforations were then made in the tissue in accordance with the 35 original pattern.

The sheet was allowed to dry overnight, removed, cut square, and labelled for identification.

CLAIMS:

1. A unitary homolaminar tissue graft construct formed from multiple strips of intestinal submucosa tissue, said strips of intestinal submucosa tissue delaminated from both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, said construct consisting essentially of submucosal tissue wherein said construct has a greater surface area than any one of the individual strips used to form said construct.

2. The tissue graft construct of claim 1, wherein the strips of intestinal submucosa tissue consist essentially of the tunica submucosa, the muscularis mucosa and the stratum compactum of the tunica mucosa.

3. The tissue graft construct of claim 1, wherein the strips of intestinal submucosa tissue are conditioned to have a strain of no more than 20%.

4. A method for forming a unitary homolaminar tissue graft construct from multiple strips of submucosal tissue delaminated from both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, wherein said construct has a greater surface area than any one of the individual strips used to form said construct said method comprising the steps of

25 forming a first layer of submucosal tissue by locating strips of submucosal tissue on a first surface wherein the strips of submucosal tissue of the first layer are adjacent to one another and in contact with one another without substantial overlap between one another;

30 overlapping the first layer of submucosal tissue with a second layer of submucosal tissue strips wherein the strips of submucosal tissue of the second layer are adjacent to one another and in contact with one another without substantial overlap between one another and the strips of submucosal tissue of the second layer are offset

in relationship to the submucosal strips of the first layer so that the contacting edges of the individual strips of submucosal tissue of the first layer are bridged by the strips of submucosal tissue of the second layer; and

5 compressing the overlapped portions of the strips of submucosal tissue between the first surface and a second surface under conditions allowing at least partial dehydration of the compressed submucosal tissue.

10 5. The method of claim 4, further comprising the step of applying a vacuum to the submucosal tissue during compression thereof.

6. The method of claim 5, wherein the applied vacuum coacts with the two surfaces to compress the submucosal tissue.

15 7. The method of claim 4, further comprising the step of applying heat to the submucosal tissue during compression thereof.

8. The method of claim 4, wherein at least one of the surfaces is water permeable.

20 9. The method of claim 4, wherein the two surfaces are complementary nonplanar surfaces wherein compression between the two surfaces produces a tissue graft construct shaped to conform to those surfaces.

25 10. A unitary homolaminar tissue graft construct, consisting essentially of intestinal submucosal tissue delaminated from both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, said homolaminar construct formed to have pseudoisotropic properties.

30 11. The tissue graft construct of claim 10, wherein the intestinal submucosa tissue consist essentially of the tunica submucosa, the muscularis mucosa and the stratum compactum of the tunica mucosa.

12. The tissue graft construct of claim 10, wherein the intestinal submucosa tissue is conditioned to have a strain of no more than 20%.

5 13. A method of preparing a multi-laminate tissue graft construct having pseudoisotrophic properties, said construct formed from at least three strips of intestinal submucosal tissue delaminated from both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, each strip of 10 intestinal submucosal tissue having a longitudinal axis corresponding to the predominant orientation of the collagen fibers in the submucosal tissue strips, said method comprising the steps of

15 overlaying said first strip with at least two additional strips of submucosal tissue so that the longitudinal axis of each individual strip of submucosal tissue forms an angle of about $180^\circ/N$ with the longitudinal axes of at least two other strips of overlaid submucosal tissue, wherein $N =$ the total number of strips of 20 submucosal tissue;

compressing at least the overlapped portions of the strips of submucosal tissue between two surfaces under conditions allowing at least partial dehydration of the compressed submucosal tissue.

25 14. The method of claim 13, wherein at least one of the surfaces is water permeable.

30 15. A method of preparing a multi-laminate tissue graft construct having pseudoisotrophic properties, said construct formed from multiple sheets of intestinal submucosal tissue, wherein each of said sheets comprise overlapped strips of submucosal tissue delaminated from both the tunica muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, each of said strips having a longitudinal axis corresponding to the predominant orientation of the collagen fibers in the 35

submucosal tissue strips, wherein the longitudinal axes of the strips of submucosal tissue of each sheet are substantially parallel to one another, said method comprising the steps of

5 overlaying said first sheet of submucosal tissue with at least two additional sheets of submucosal tissue so that the longitudinal axes of the strips of submucosal tissue of one sheet form an angle of about $180^\circ/S$, with the longitudinal axes of the strips of submucosal tissue of at least two of the overlaid sheets of submucosal tissue, wherein $S =$ the total number of sheets of submucosal tissue; and

10 compressing the sheets of submucosal tissue between two surfaces under conditions allowing at least partial dehydration of the compressed submucosal tissue.

15 16. The method of claim 15, further comprising the step of applying a vacuum to the submucosal tissue during compression thereof.

20 17. The method of claim 16, wherein the applied vacuum coacts with the two surfaces to compress the submucosal tissue.

25 18. The method of claim 15, further comprising the step of applying heat to the submucosal tissue during compression thereof.

30 19. The method of claim 15, wherein at least one of the surfaces is water permeable.

35 20. The method of claim 15, wherein the two surfaces are complementary nonplanar surfaces wherein compression between the two surfaces produces a tissue graft construct shaped to conform to those surfaces.

21. A multi-laminate tissue graft construct having pseudoisotrophic properties, said construct formed from multiple sheets of intestinal submucosal tissue, wherein each of said sheets comprise overlapped strips of submucosal tissue delaminated from both the tunica

muscularis and the luminal portion of the tunica mucosa of a warm blooded vertebrate, each of said strips having a longitudinal axis corresponding to the predominant orientation of the collagen fibers in the submucosal tissue strips, wherein the longitudinal axes of the strips of submucosal tissue of each respective sheet are substantially parallel to one another

5 22. The multi-laminate tissue graft construct of claim 21, wherein the overlapped strips of submucosal tissue comprise strips of submucosal tissue each having a portion overlapped by a portion of an adjacent strip of submucosal tissue.

10 23. The multi-laminate tissue graft construct of claim 21, wherein the overlapped strips of submucosal tissue comprise a first layer of submucosal tissue strips overlaid with a second layer of submucosal tissue strips, wherein the strips of submucosal tissue of each respective layer are positioned adjacent to and in contact with one another without substantial overlap between the adjacent 15 strips of submucosal tissue, the strips of submucosal tissue of said second layer being offset in relationship to the submucosal strips of said first layer so that the contacting edges of the individual strips of submucosal tissue of the first layer are bridged by the strips of submucosal tissue of the second layer.

20 24. The graft of claim 21 wherein the strips of submucosal tissue consist essentially of the tunica submucosa, the muscularis mucosa and the stratum compactum of the tunica mucosa.

25 25. The graft of claim 21 wherein the strips of intestinal submucosa tissue are conditioned to have a strain of no more than 20%.

30 26. The tissue graft material of claim 21, wherein the segment of intestinal tissue comprises the tunica submucosa and basilar tissue of the tunica mucosa.

1/2

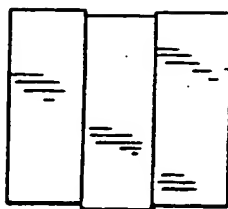


FIG. 1a

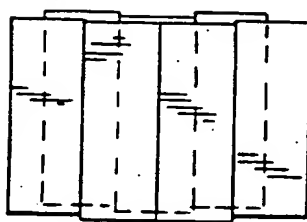


FIG. 1b

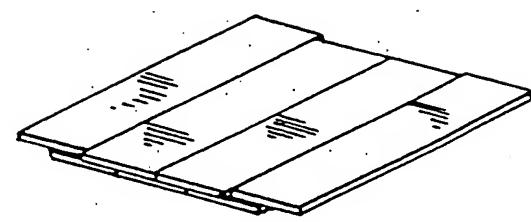


FIG. 1c



FIG. 2a

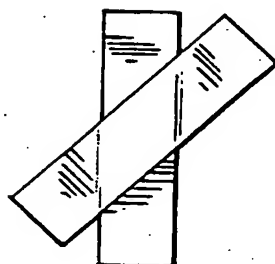


FIG. 2b

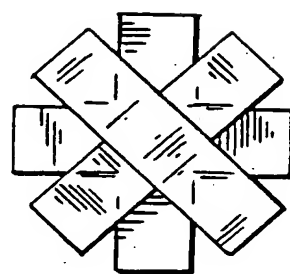


FIG. 2c

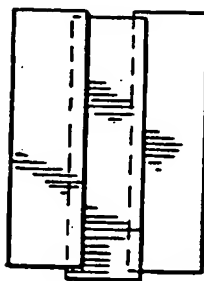


FIG. 3a

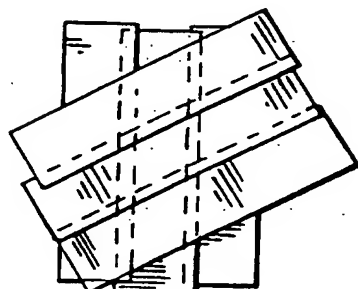


FIG. 3b

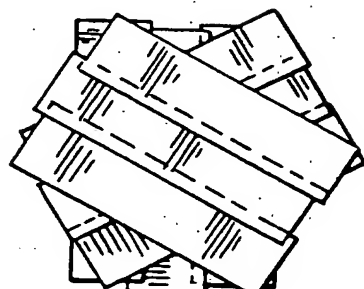
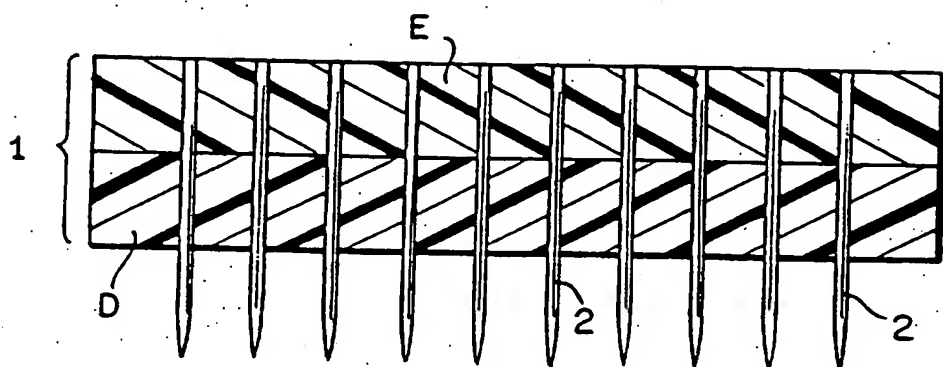
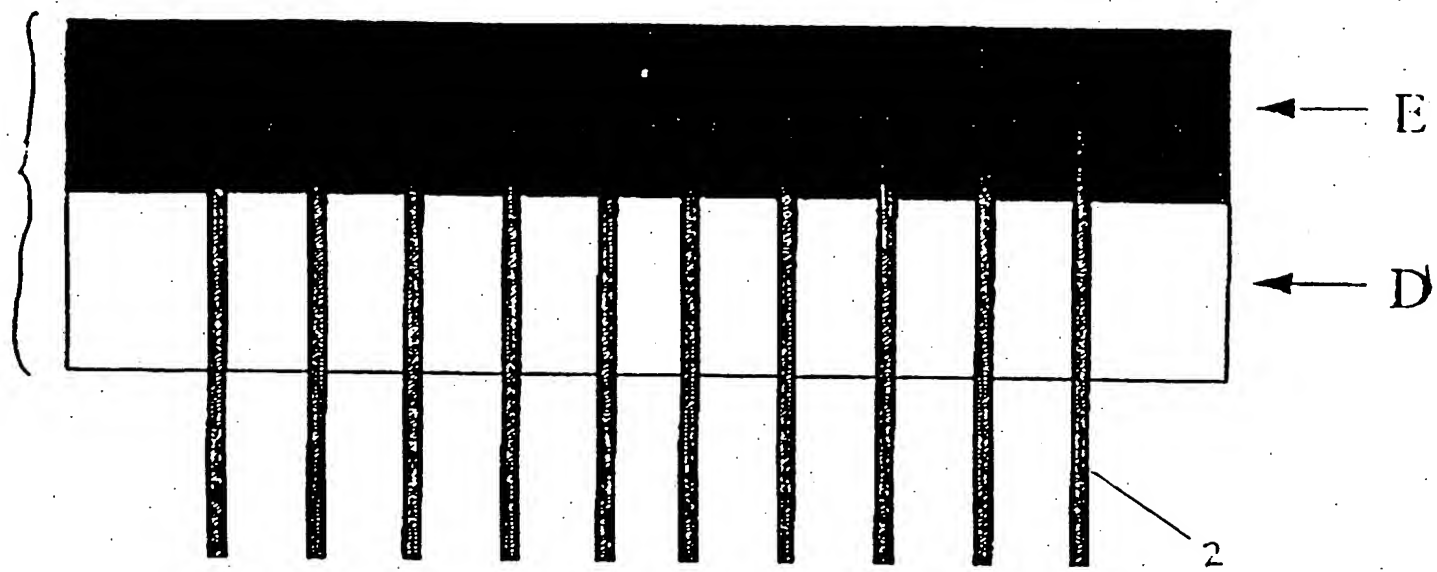


FIG. 3c

FIG. 4





INTERNATIONAL SEARCH REPORT

International application No.
PCT/US96/04721

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : A61K 35/38; A61F 2/02

US CL : 424/551; 623/11

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 424/551; 623/11

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

APS, CA, BIOSIS, MEDLINE

search terms: submucosa, graft

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	US 3,562,820 A (BRAUN) 16 February 1971 (16.02.71), column 2, lines 47-64.	1, 3

Further documents are listed in the continuation of Box C. See patent family annex.

Special categories of cited documents:	"T"	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
A document defining the general state of the art which is not considered to be of particular relevance	"X"	document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
E earlier document published on or after the international filing date	"Y"	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
L document which may throw doubt on priority (claims) or which is cited to establish the publication date of another citation or other special reason (as specified)	"&"	document member of the same patent family
O document referring to an oral disclosure, use, exhibition or other means		
P document published prior to the international filing date but later than the priority date claimed		

Date of the actual completion of the international search

22 JULY 1996

Date of mailing of the international search report

29 JUL 1996

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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US96/04721

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be examined, the appropriate additional examination fees must be paid.

Group I, claims 1-12, drawn to a homolaminar tissue graft and method of making same.

Group II, claims 13-26, drawn to a multilaminate tissue graft and method of making same.

The inventions listed as Groups I and II do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: The tissue grafts of Groups I and II do not share a special technical feature as they are different compositions made by two different processes. The second method is cited in Group II as opposed to adding a third group, because there is no additional burden of search.

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